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A local human V δ 1 T cell population is associated with survival in nonsmall-cell lung cancer

Yin Wu[®]^{1,2,3,4,55}[∞], Dhruva Biswas[®]^{1,2,5,55}, leva Usaite¹, Mihaela Angelova[®]², Stefan Boeing⁶, Takahiro Karasaki[®]^{1,2}, Selvaraju Veeriah¹, Justyna Czyzewska-Khan¹, Cienne Morton³, Magdalene Joseph^{3,4}, Sonya Hessey^{1,7}, James Reading[®]^{1,8}, Andrew Georgiou^{1,8}, Maise Al-Bakir², TRACERx Consortium^{*}, Nicholas McGranahan^{1,9}, Mariam Jamal-Hanjani[®]^{1,7}, Allan Hackshaw[®]¹⁰, Sergio A. Quezada[®]^{1,8}, Adrian C. Hayday[®]^{3,4}[∞] and Charles Swanton[®]^{1,2}[∞]

Murine tissues harbor signature $\gamma\delta$ T cell compartments with profound yet differential impacts on carcinogenesis. Conversely, human tissue-resident $\gamma\delta$ cells are less well defined. In the present study, we show that human lung tissues harbor a resident V $\delta1$ $\gamma\delta$ T cell population. Moreover, we demonstrate that V $\delta1$ T cells with resident memory and effector memory phenotypes were enriched in lung tumors compared with nontumor lung tissues. Intratumoral V $\delta1$ T cells possessed stem-like features and were skewed toward cytolysis and helper T cell type 1 function, akin to intratumoral natural killer and CD8⁺ T cells considered beneficial to the patient. Indeed, ongoing remission post-surgery was significantly associated with the numbers of CD45RA⁻CD27⁻ effector memory V $\delta1$ T cells in tumors and, most strikingly, with the numbers of CD103⁺ tissue-resident V $\delta1$ T cells in nonmalignant lung tissues. Our findings offer basic insights into human body surface immunology that collectively support integrating V $\delta1$ T cell biology into immunotherapeutic strategies for nonsmall cell lung cancer.

mmune checkpoint inhibitors (CPIs) have revolutionized the treatment of cancer by providing durable remissions, albeit for a minority of patients. CPIs work, at least in part, by de-repressing tumor (neo)antigen-specific $\alpha\beta$ T cells. Thus, many efforts to improve efficacy have focused on this axis^{1,2}. Indeed, some utility of tumor mutational burden (TMB)^{3,4}, presence and quality of CD8⁺ T cells^{5,6} and major histocompatibility complex (MHC) class I loss of heterozygosity⁷ in predicting responses to CPIs provide collective evidence of the contributions of antigen-specific $\alpha\beta$ T cells. Nevertheless, a high TMB does not guarantee responses to CPIs and a lack of MHC-I and/or low TMB does not preclude good responses^{8,9}.

Alongside $\alpha\beta$ T cells, we and others have previously demonstrated that $\gamma\delta$ T cells are also found within tumors¹⁰⁻¹⁴. These cells are evolutionarily conserved, implying a vital and nonredundant role and, although they are often less abundant than $\alpha\beta$ T cells, this disparity may be compensated by their polyclonal response potentials compared with the highly clonotypic responses of $\alpha\beta$ T cells¹⁵. Indeed, it is well established that $\gamma\delta$ T cells protect against cancer in mice, both independently and synergistically with $\alpha\beta$ T cells^{16,17}, highlighting the need to better assess their importance in human cancers^{18,19}. Although such studies have been limited by the availability of technologies to rigorously identify, isolate and examine $\gamma\delta$ T cells, a recent in silico study of >5,000 cancer patients demonstrated that, of all 22 immune cell types studied, the intratumoral $\gamma\delta$ T cell signature was the trait most significantly associated with remission^{10}.

Similar to $\alpha\beta$ T cells, $\gamma\delta$ T cells are also composed of distinct subsets occupying different functional niches. It is clear in the murine setting that body surface tissues in which carcinomas arise harbor signature tissue-specific $\gamma\delta$ T cell subsets²⁰. We recently demonstrated that human breast epithelium is enriched in the V δ 1 subset of y8 T cells and that these cells possess potent anti-tumor functions, including the capacity to kill transformed cells in vitro and produce tumor-antagonistic cytokines, for example, interferon-y (IFN- γ)¹¹. Moreover, activation of these V δ 1 T cells did not require cognate peptide-MHC but instead the cells responded to conserved signals of tissue stress, via innate receptors, entirely distinct from co-located $\alpha\beta$ T cells¹¹. In other words, V δ 1 T cells were an independent, but potentially synergistic, population of anti-tumor lymphocytes in the tumor microenvironment (TME). Furthermore, in patients with aggressive triple-negative breast cancers (TNBCs), the intratumoral presence of Vδ1 T cells was more significantly associated with survival than co-located $\alpha\beta$ T cells. A subsequent study employing CIBERSORT demonstrated that intratumoral γδ T cells also predicted survival in a larger cohort of 169 patients with TNBCs in the METABRIC dataset²¹. Recent studies have described similar populations of resident V\delta1 T cells in colorectal cancers, where their presence was associated with lower-stage disease¹², and in hepatocellular carcinomas, where the presence of total $\gamma\delta$

¹Cancer Research UK Lung Cancer Centre of Excellence, University College London Cancer Institute, London, UK. ²Cancer Evolution and Genome Instability Laboratory, The Francis Crick Institute, London, UK. ³Peter Gorer Department of Immunobiology, School of Immunology & Microbial Sciences, King's College London, London, UK. ⁴Immunosurveillance Laboratory, The Francis Crick Institute, London, UK. ⁵Bill Lyons Informatics Centre, University College London Cancer Institute, London, UK. ⁶Bioinformatics & Biostatistics and Software Development & Machine Learning Team, The Francis Crick Institute, London, UK. ⁷Cancer Metastasis Lab, University College London Cancer Institute, London, UK. ⁸Cancer Immunology Unit, Research Department of Haematology, University College London Cancer Institute, London, UK. ⁹Cancer Genome Evolution Research Group, University College London Cancer Institute, London, UK. ¹⁰Cancer Research UK & University College London Cancer Trials Centre, University College London, UK. ⁵⁵These authors contributed equally: Yin Wu, Dhruva Biswas. *A list of authors and their affiliations appears at the end of the paper. ⁵⁵e-mail: yin.wu@kcl.ac.uk; adrian.hayday@kcl.ac.uk; charles.swanton@crick.ac.uk T cells was associated with survival¹⁴. Thus, we hypothesize that human body surface tissues, similar to their murine counterparts, harbor signature tissue-resident $\gamma\delta$ T cells contributing to cancer immunosurveillance.

To test this hypothesis in human lung, we have leveraged samples and clinical data collected from the TRACERx (Tracking non-small-cell lung Cancer Evolution through therapy (Rx)) Study²². Nonsmall cell lung cancers (NSCLCs) are cancers of unmet clinical need. Although outcomes are better for patients with early-stage disease, overall outcome is poor with a 5-year survival rate of <20%²³. These cancers are clearly susceptible to immunosurveillance, as witnessed by instances of successful CPI therapy and evidence of immune editing in treatment-naive primary lung cancers²⁴. However, despite harboring high TMBs²⁵, only a minority of patients with NSCLCs respond to CPI therapy. By employing flow cytometry, quantitative T cell receptor-sequencing (TCR-seq) and RNA-sequencing (RNA-seq), we now find that both nontumor (NT) human lung tissues and NSCLCs harbor resident populations of $\gamma\delta$ T cells, particularly enriched in the V δ 1 subtype. Moreover, these Vδ1 T cells possess a T-cytolytic type 1 (Tc1) phenotype that is well established as beneficial to patients with cancer²⁶⁻²⁸. Finally, the presence of V δ 1 T cells in both NT tissue and lung tumors was significantly associated with remission. The former association is of particular interest because cell populations in NT tissue, as opposed to tumors, may be retained in situ postoperatively.

Results

Vol T cells are present in lung epithelium and enriched in NSCLCs. To characterize the T cell landscape in human lung epithelium and NSCLCs, we examined tissue and tumor-infiltrating lymphocytes (TILs) isolated from NT lung tissues and paired tumors of patients with surgically resected NSCLCs collected from the TRACERx Study (Supplementary Table 1). NT tissues were taken as far as possible from tumors at primary surgery and hematoxylin and eosin (H&E)-stained sections examined afterwards by an accredited histopathologist to ensure that samples were tumor free. TILs were isolated by enzymatic digestion and cryopreserved before thawing for use in downstream assays without further manipulation or expansion (see Methods). TILs were immunophenotyped by flow cytometry and absolute T cell counts established using quantitative TCR-seq of region-matched genomic (g)DNA extracted from bulk tissues and tumors (Fig. 1a). Samples were chosen based on the availability of banked TILs and region-matched bulk DNA from patients with at least one follow-up visit after surgical resection (see Methods). Where available, paired contemporaneous peripheral blood mononuclear cells (PBMCs) were also immunophenotyped by flow cytometry to contextualize findings with the well-characterized blood yo T cell compartment. No other selection criteria were applied.

Similar to their presence in peripheral blood (median, interguartile range (IQR): 2.49%, 0.80–6.76%), $\gamma\delta$ T cells detected by flow cytometry comprised a small fraction of total T cells in both NT lung tissues (2.22%, 1.39-5.26%) and tumors (1.15%, 0.85-3.05%), with considerable interindividual variation (Fig. 1b). In contrast to peripheral blood, where most $\gamma\delta$ T cells expressed V δ 2, most $\gamma\delta$ T cells found in lung tissues and tumors expressed Vo1 (47.6%, 30.8-74.4% and 46.8%, 32.1-70.0% respectively), consistent with a well-established enrichment of Vδ1 T cells in other tissues²⁹ (Fig. 1b and Extended Data Fig. 1a,b). Complementary to flow cytometry, we employed quantitative TCR-seq of gDNA at the TCR α/δ (TRA/ TRD) locus from matched NT tissues and tumor regions as an independent assay of $\alpha\beta$ and $\gamma\delta$ T cells, and to assess their absolute numbers per unit of tissue (Fig. 1c). Indeed, there was a strong and significant correlation between the proportion of major V δ subsets $(V\delta1 and V\delta2)$ detected by flow cytometry and TCR-seq (Extended Data Fig. 1c). We then mapped flow phenotyping proportions

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(that is, proportion of CD3⁺ T cells positive for CD4⁺/CD8⁺) on to absolute TCR-seq counts of T cells to derive more granular T cell subset numbers (Fig. 1c). Consistent with flow cytometry, TCR-seq also revealed that $\gamma\delta$ T cells form a minority subset of total T cells in NT lung tissues (2.70%, 1.30-6.23%) and tumors (1.68%, 0.97-2.60%), and that the V δ 1 subset comprises most of these cells in both NT lung tissues (54.2%, 35.9-76.7%) and tumors (62.0%, 29.5-79.6%) (Fig. 1c). Although NT lung tissues were macroscopically and microscopically tumor free, it was not practical to obtain normal lung tissue from healthy donors to exclude potential effects of tumors on the immune microenvironment of NT lung tissues. To address this issue, we utilized data generated by the Genotype-Tissue Expression (GTEx) project³⁰, a comprehensive public resource of tissue-specific gene expression from 54 nondiseased tissue sites across almost 1,000 individuals. We found that TRDC, a gene expressed by all γδ T cells, was highly expressed in lung tissues compared with other tissue sites (Extended Data Fig. 2a). Furthermore, expression of TRDV1 (Vδ1 T cells) was higher than TRDV2 (Vδ2 T cells) within lung tissue (Extended Data Fig. 2b,c), consistent with our own data. Thus, we conclude that $\gamma\delta$ T cells, particularly the Vol subset, are present in nonmalignant lung tissue at a steady state.

In many tissues, CD103 has been adopted as a marker of tissue-resident memory (T_{RM}) status, absent on most peripheral blood T cells³¹. Consistent with previous studies, NT lung tissues harbored a CD103⁺CD8⁺TCR $\alpha\beta$ T_{RM} population (median, IQR: 35.3%, 25.9–55.0%) (Fig. 2a,b)^{32,33}. Similar to the CD8⁺ T cell compartment, we found that many, albeit not all, V δ 1 T cells in NT lung tissues displayed a CD103⁺ T_{RM} phenotype, particularly in some patients (18.8%, 3.35–62.4%) (Fig. 2a,b). In contrast, most CD4⁺ and V δ 2 T cells in NT lung tissues were CD103⁻ as were their counterparts in peripheral blood (Fig. 2a,b). Thus, the resident CD103⁺ T cell compartment in NT lung tissues mostly comprised CD8⁺ and V δ 1 T cells.

Compared with NT lung tissues, tumors harbored a greater proportion of CD8⁺ (median, IQR: 65.6%, 57.7–75.3%), V δ 1 (66.6%, 34.4–78.5%) and V δ 2 T cells (10.3%, 5.03–18.2%) expressing CD103 whereas the proportion of CD103 expressing CD4⁺ T cells was reduced (Fig. 2b). By mapping flow phenotyping proportions on to TCR-seq counts, we observed significantly greater absolute numbers of CD8⁺ and V δ 1 T cells with the CD103 phenotype in tumors compared with NT lung tissues; conversely, this was not so for CD4⁺ and V δ 2 T cells (Fig. 2c).

Intratumoral Vδ1 T cells possess a Tc1 functional phenotype in situ. Intratumoral CD8⁺ T_{RM} cells have been associated with survival in several cancers^{34,35} including NSCLCs³⁶. However, the potential contributions of human Vδ1 T cells are less well described and more contentious³⁷. Similar to CD8⁺ αβ T cells, γδ T cells have been operationally classified into effector T cell/memory T cell subsets associated with defined effector functions based on CD45RA and CD27 expression³⁸. When we compared Vδ1 T cells in tumors versus NT tissues, we observed an overt shift toward a CD45RA⁻CD27⁻ effector memory T cell (T_{EM}) phenotype by flow cytometry (Fig. 3a,b). Similar to their CD8⁺ counterparts, Vδ1 T_{EM} cells have been associated with helper type 1 T cell (T_H1 cell) cytokine production (IFN-γ) and cytolysis³⁸, but whether intratumoral Vδ1 T cells possess this patient-beneficial Tc1 phenotype^{27,28} in situ has remained unclear.

To address this, we sorted bulk V δ 1 T cells from disaggregated NT tissues and lung tumors (n=2 and n=5 patients, respectively) from which there was sufficient material. For comparison, we also sorted, from NT lung tissues and tumors, bulk V δ 2 (n=3 and n=3 patients), CD4⁺ (n=8 and n=9 patients), CD8⁺ (n=7 and n=9), and regulatory T cells (T_{reg} cells) (n=1 and n=5 patients) and natural killer (NK) cells (n=8 and n=7 patients) (Extended Data Fig. 3a). Disaggregated TILs were immediately frozen and

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Fig. 1 Experimental design and \gamma\delta T cell composition in lung tissues and NSCLCs. a, Overview of study design. Paired tumor regions (red) and NT lung tissues (blue) collected under the TRACERx Study were enzymatically digested to extract tissue/TILs. TILs were cryopreserved and thawed at a later date for flow cytometry ± RNA-seq. In parallel, gDNA was extracted from undigested matched tumor regions and NT lung tissues and sent for subsequent quantitative TCR-seq. In addition, PBMCs were isolated from contemporaneous blood draws and cryopreserved before subsequent thaw for flow cytometry. b, Percentage of CD3⁺ T cells staining for TCR $\gamma\delta$ (left) and percentage of TCR $\gamma\delta$ T cells staining for V δ 1 (middle) and V δ 2 (right) in PBMCs (blood), NT lung tissues (tissue) and tumors (tumor). Not all patients had paired samples. The bar represents the median. The Kruskal-Wallis test with post-hoc Dunn's test corrected for multiple testing was used. **c**, Absolute counts of total T cells, $\alpha\beta$ T cells (TRA), $\gamma\delta$ T cells (TRD) and V δ 1 (*TRDV1*) and V δ 2 (*TRDV2*) T cells per microgram of DNA determined by TCR-seq. Absolute counts of CD4⁺ $\alpha\beta$ T cells (CD4) and CD8⁺ $\alpha\beta$ T cells (CD8) were determined by mapping the proportion of CD3⁺/TCR $\gamma\delta^-$ T cells staining for CD4 or CD8 in flow cytometry analysis of paired TILs. No significant differences were observed within demarcated T cell subsets between NT tissues and tumors. Samples with <1 cell μg^{-1} of DNA were not plotted for the purposes of visualization. The bar represents the median. A two-tailed Mann-Whitney *U*-test was used within demarcated T cell subsets. Significant *P* values are shown. NS, not significant. The *n* numbers and datapoints represent independent patients.

subsequently thawed, stained and sorted at 4°C directly into lysis buffer for RNA-seq, thereby maximizing preservation of the cells' in situ transcriptomes.

Cell types clustered together in a principal component analysis (PCA) of the 500 most variably expressed genes (Extended Data Fig. 3b) and expressed the anticipated canonical lineage markers (Extended Data Fig. 3b). Notably, V δ 1 T cells clustered together with CD8⁺ T cells and NK cells in this unsupervised analysis (Extended Data Fig. 3b). To determine the functional skew of V δ 1 T cells, we restricted our analysis to canonical transcription factors and effector molecules associated with T cell function and found

that intratumoral V δ 1 T cells expressed the T_H1 cell-specific master transcription factor Tbet (*TBX21*) as well as transcripts for granzymes (*GZMs*), perforin (*PRF1*) and IFN- γ (*IFNG*) to levels comparable with those of CD8⁺ T cells and NK cells (Fig. 3c). Moreover, intratumoral V δ 1 T cells did not express T_H17 cell-associated genes or the T_H17⁻ cell-specific master transcription factor ROR γ t (*RORC*). By contrast, these genes were expressed to a variable degree by co-located CD4⁺ T cells (Fig. 3c). To contextualize the in situ function of intratumoral V δ 1 T cells, we conducted a PCA using these restricted genes of interest expressed by intratumoral V δ 1, V δ 2, CD4⁺, CD8⁺ and T_{reg} cells and NK cells. The PCA demonstrated that intratumoral V δ 1 T cells transciptomically resemble CD8⁺ T cells and NK cells in function (Fig. 3d). To further contextualize our results, we reanalyzed expression of these genes in comparable cell types isolated from the peripheral blood of healthy volunteers as part of the Blood Atlas Project³⁹ (Extended Data Fig. 4). Intratumoral V δ 1 T cells resemble peripheral blood $\gamma\delta$ T cells in the expression of genes associated with T_H1 cell, T_H2 cell, cytolytic and inhibitory functions. Notably, peripheral blood $\gamma\delta$ T cells showed some evidence for T_H17 cell-associated gene expression (Extended Data Fig. 4), whereas this was not seen for intratumoral V δ 1 T cells (Fig. 3c). Most probably, T_H17 cell-associated gene expression in blood reflected the predominance of V δ 2 T cells that have been reported to produce interleukin (IL)-17, albeit rarely⁴⁰⁻⁴².

Next, we validated the functional potential of intratumoral V δ 1 T cells in vitro by stimulation of TILs with phorbol 12-myristate 13-acetate (PMA) and ionomycin, which mimics TCR signaling. Stimulated TILs were then stained for surface lineage markers and CD107A, a marker of cytotoxic degranulation, as well as for intracellular cytokines. Consistent with their gene expression profile, V δ 1 T cells produced IFN- γ and degranulated on activation (Fig. 3e and Extended Data Fig. 5). Moreover, we could find no evidence of IL-17A production by these cells in contrast to co-located CD4⁺ T cells (Fig. 3e and Extended Data Fig. 5).

Previous studies have demonstrated that tissue-associated V\delta1 T cells may also be activated by the innate NKG2D receptor without requirement for contemporaneous TCR signaling^{11,12,43}. Consistent with an innate, non-TCR/nonclonotypic response mode, we observed no significant clonal focusing of the TCRS chain in Vo1 T cells in tumors compared with NT tissues (Extended Data Fig. 6a,b). The gDNA-based TCRy (TRG) sequencing from bulk tissues is inherently problematic because $\alpha\beta$ T cells often harbor productive rearrangements of TRG genes⁴⁴. Nevertheless, RNA-seq of sorted V\delta1 T cells from tumors, albeit in a limited cohort, demonstrated a diverse expression of Vy chains in most patients, in contrast to V82 T cells which predominantly employed Vy9 (Extended Data Fig. 6c), further supporting a nonclonal innate response mode for Vô1 T cells. Moreover, we also observed a positive and significant correlation of the presence of intratumoral V δ 1 T_{EM} cells with region-matched intratumoral expression of transcripts for NKG2D ligands (Fig. 3f).

Intratumoral Vδ1 T cells demonstrate features of tissue residency and stemness. Whereas CD103 expression is often associated with T_{RM} status in T cells⁴⁵, this is not always the case^{46,47}. Seminal studies by independent teams have collectively established a core transcriptional profile of T_{RM} cells. Specifically, T_{RM} cells upregulate genes associated with tissue retention and homing (*CD69*, *CXCR6*, *ITGAE* and *ITGA1*) and downregulate genes associated with tissue egress (*CCR7*, *S1PR1* and *SELL*) compared with circulating T cells^{48,49}. Thus, we compared the expression of these genes in intratumoral Vδ1 and CD8⁺ T cells, which were predominantly CD103⁺, versus intratumoral CD4⁺ T cells, which mostly lacked CD103 (Fig. 2b), and found that this tissue-resident profile was shared by the Vδ1 and CD8⁺ T cells, consistent with a bona fide $\rm T_{\rm RM}$ status (Fig. 3g and Extended Data Fig. 7).

Recent studies have identified a distinct subset of stem-like memory CD8⁺ T cells in chronic infection and cancer defined by their expression of the transcription factors/regulators *EOMES*, *TCF7* and *TOX* and the IL-7 receptor (*IL7R*)⁵⁰⁻⁵². These cells retain the capacity to proliferate despite chronic inflammatory stimuli and can give rise to highly functional effector cells implicated in tumor control and responses to programmed cell death protein 1 (PD-1) blockade^{50,53}. Intratumoral Võ1 T cells resembled stem-like CD8⁺ T cells in their expression of *EOMES*, *TCF7* and *TOX* distinct from intratumoral CD4⁺ T cells (Fig. 3h). Intratumoral Võ1 T cells also expressed *IL7R* (Fig. 3h), albeit at a somewhat lower level, possibly because these cells are maintained by the epithelial-associated cytokine, IL-15 (ref. ⁵⁴).

Presence of Vo1 T cells predicts ongoing remission in resected NSCLCs. Given that V\delta1 T_{RM} and T_{EM} cells were enriched in tumors relative to NT lung tissues (see above), intratumoral V\delta1 T cells possess a Tc1 phenotype and these cells resemble stem-like CD8+ T cells, we examined their status in relation to clinical outcome in our cohort. Three patients who had incompletely excised primary tumors were excluded from this outcome analysis (Supplementary Table 1). To assess absolute numbers of T_{RM} (CD103⁺) and T_{EM} (CD45RA-CD27-) cells per unit of tissue/tumor, we mapped flow phenotyping proportions on to TCR-seq counts. Within tumors, the presence of V\delta1 $T_{\mbox{\tiny EM}}$ cells was significantly associated with increased relapse-free survival (RFS) whereas this was only a trend for co-located V82, CD4⁺ and CD8⁺ T_{EM} cells (Fig. 4a). Importantly, the association of intratumoral V\delta1 $T_{\mbox{\tiny EM}}$ cells with improved RFS was not simply a reflection of less advanced disease because we found more intratumoral V\delta1 $T_{\mbox{\tiny EM}}$ cells in advanced stages (Extended Data Fig. 8a). Furthermore, we found no association of intratumoral Vδ1 T_{EM} cells with primary tumor size, age, histology or smoking status (Extended Data Fig. 8a). The presence of V δ 1 T_{RM} cells in tumors trended toward association with increased RFS (Extended Data Fig. 8b). Conspicuously, we found the presence of V δ 1 T_{RM} cells in NT tissue to be highly and significantly associated with improved RFS, consistent with an epithelial immunosurveillance role proposed for these cells¹⁶ (Fig. 4b). There was no difference in the number of Vδ1 T_{RM} cells in NT tissue with regard to stage of disease, size of primary, age, histology or smoking status (Extended Data Fig. 8c).

Given the association of Võ1 T cells in both NT tissues and tumors with ongoing remission, we asked whether such an association might exist for the delta (Δ) of the absolute numbers of Võ1 T_{EM} cells and Võ1 T_{RM} cells in NT tissues and paired tumors. No clear association of Δ Võ1 T_{EM} or Δ Võ1 T_{RM} cells with clinical outcome was observed (Extended Data Fig. 8d), suggesting that each subset, that is, intratumoral Võ1 T_{EM} cells and NT tissue Võ1 T_{RM} cells, has beneficial impacts independent of the status of their counterpart cells in the reciprocal tissue sites.

To explore further the contributions of tissue-resident V δ 1 T cells, we used the nucleotide sequence of the V δ 1 complementarity -determining region 3 (CDR3) as a molecular fingerprint to track

Fig. 2 | NT lung tissue harbors tissue-resident Vô1T cells that are enriched in NSCLCs. a, Representative plots of CD103 expression (%) by flow cytometry on Vô1 (representing n = 20, n = 23 and n = 22 patients for blood, NT tissue and tumor, respectively) and Vô2 (representing n = 20, n = 23 and n = 21 patients for blood, NT tissue and tumor, respectively), CD4+ (representing n = 20, n = 24 and n = 24 patients for blood, NT tissue and tumor, respectively) and CD8+ T cells (representing n = 20, n = 24 and n = 24 patients for blood, NT tissue and tumor, respectively) and CD8+ T cells (representing n = 20, n = 24 and n = 24 patients for blood, NT tissue and tumor, respectively) and CD8+ T cells (representing n = 20, n = 24 and n = 24 patients for blood, NT tissue and tumor, respectively) were isolated from blood, NT tissue and tumor of one patient. **b**, Summary flow cytometry data of CD103 expression in T cell subsets isolated from blood (Bld), NT tissues (Tis) and tumors (Tum). Not all patients had paired samples. The bar represents the median. A Kruskal-Wallis test with a post-hoc Dunn's test corrected for multiple testing was used. **c**, Absolute counts of CD103+ T_{RM} CD4+, CD8+, Vô1 and Vô2 T cells per microgram of DNA from NT tissues and tumors. Samples with <1 cell μg^{-1} of DNA were not plotted for the purposes of visualization. Not all patients had paired samples. The bar represents the median. A two-tailed Mann-Whitney *U*-test was used within demarcated T cell subsets. Significant *P* values are shown. NS, not significant. The *n* numbers and datapoints represent independent patients.

unique V δ 1 T cell clones between NT tissues and paired tumors. On average, approximately a quarter of unique V δ 1 CDR3 sequences present in tumors were found in paired NT tissues (median,

IQR: 27.3%, 16.5–35.4%) (Fig. 4c). By contrast, fewer unique TCR α clones present in tumors were also found in paired NT tissues (15.1%, 8.68–19.2%) (Fig. 4d), despite the lower potential for



diversity in TCR α compared with TCR δ . Indeed, these data are consistent with a greater proportion of intratumoral $\alpha\beta$ T cells (predominantly CD4⁺) being derived from peripheral blood as opposed to the tissue-resident pool that is the probable source of intratumoral V δ 1 T cells (Fig. 2b). When examined in relation to clinical outcome, patients with a greater proportion of intratumoral V δ 1 T cells also found in paired NT tissues were more likely to remain in remission, consistent with the cells' proposed immunosurveillance function in steady-state tissues (Fig. 4c). Again, this was not the case for the $\alpha\beta$ T cell compartment (Fig. 4d).

Acknowledging the limited size of our cohort, we sought to validate our findings in a larger public dataset. The TCR δ locus is excised during TCR α rearrangement in $\alpha\beta$ T cells⁵⁵. Thus, we used the expression of *TRDV1* transcripts as a proxy for V δ 1 T cells and *TRDC* transcripts as a proxy for total $\gamma\delta$ T cells, thereby assessing the association of these cells with NSCLC survival in The Cancer Genome Atlas (TCGA). Consistent with observations in our own cohort, when split on median expression, we found a significant association of high *TRDV1* expression with favorable overall survival (OS; Fig. 5a), although there was no association with *TRDC* expression (Fig. 5b). Moreover, the improved hazard ratio (HR) in patients with above-median *TRDV1* expression remained significant in multivariate analysis accounting for age, gender, histology, smoking status, stage and *CD4* and *CD8B* gene expression (HR=0.65, 95% confidence interval (CI)=0.48-0.88) (Fig. 5c).

Finally, to explore whether there might be a role for $V\delta1$ T cells with respect to CPI therapy responses, we reanalyzed RNA-seq data from the INSPIRE trial (NCT02644369), a phase II basket clinical trial of pembrolizumab in advanced solid cancers⁵⁶. Using *TRDV1* expression in pre-treatment tumor biopsies as a proxy for intratumoral V $\delta1$ T cells, we found that patients with above-median expression of *TRDV1* had significantly increased survival compared with those with below-median expression (Fig. 5d). Expression of *TRDC* (pan $\gamma\delta$ T cells) was also predictive, although this was probably driven by V $\delta1$ T cells because the association of *TRDV2* expression (V $\delta2$ T cells) with survival was only a trend (Extended Data Fig. 9). In addition, high expression of *CD4* (CD4⁺ T cells) did not associate with survival after pembrolizumab whereas high expression of *CD8B* (CD8⁺ T cells) trended toward association with improved survival (Extended Data Fig. 9).

Discussion

Despite recent advances in immunotherapy, NSCLCs remain a leading cause of cancer-related mortality. Although these cancers are clearly susceptible to immunosurveillance, as evidenced by instances of durable responses to CPI therapy, only a minority of patients benefit. CPIs appear to work, at least in part, by de-repressing tumor (neo)antigen-specific $\alpha\beta$ T cells. However, the body surfaces from which carcinomas arise are populated by many other immune cells. These compose organ-specific, tissue-resident immune compartments that collectively include considerably more T cells than the systemic lymphoid organs.

In mice, such compartments commonly include signature populations of tissue-resident $\gamma\delta$ T cells seeded during fetal and perinatal development²⁰. This is distinct from tissue-resident $\alpha\beta$ T cells which arrive later in development, after priming in the lymph nodes as a response to infection⁵⁷. Mice deficient in $\gamma\delta$ T cells display heightened susceptibility to de novo cancers^{16,58}, seemingly more so than mice deficient in $\alpha\beta$ T cells¹⁷, establishing a critical and nonredundant role(s) for them. Indeed, these cells appear to be an essential early source of IFN- γ^{59} , a cytokine pivotal for tumor rejection²⁶. In addition to direct effector function, human $\gamma\delta$ T cells are also capable of phagocytosis^{60,61} and professional antigen presentation to $\alpha\beta$ T cells^{60,62}. Such sentinel functions, traditionally associated with dendritic cells, are central to activating tumor (neo) antigen-specific $\alpha\beta$ T cells and to orchestrating immune responses more generally⁶³.

Despite apparently advantageous traits, $\gamma \delta$ T cell immunotherapies have failed to demonstrate convincing efficacy in solid cancers including in NSCLCs⁶⁴. However, those trials focused exclusively on the V δ 2 subset, the main subset of peripheral blood $\gamma \delta$ T cells. By contrast, our assessments of tissue-resident lymphocytes in lung tissues have added to growing evidence that human body surfaces are primarily enriched for V δ 1 $\gamma \delta$ T cells. Thus, these cells are well placed to detect malignancy vis-à-vis local/ipsilateral as well as metastatic/contralateral lung cancer recurrences, which comprise a considerable proportion of relapses after surgical resection.

Indeed, an increased presence of V δ 1 T_{RM} cells in NT lung tissues was predictive of ongoing remission in our cohort. Moreover, most V δ 1 T cells found within tumors were also of this tissue-resident phenotype, suggesting an improved capacity for tumor homing and/or retention. Strikingly, a greater proportion of shared V δ 1 T cell clones between tumors and NT tissues was significantly associated with remission in our cohort. Thus, this supports a potential patient-beneficial, cancer immunosurveillance role for human tissue-resident V δ 1 T cells that are present in steady-state NT lung tissues. Although many studies have demonstrated the prognostic utility of intratumoral TILs, these cells clearly cannot actively contribute to immunosurveillance after resection. Conversely, resident immune cells in juxtapositional normal tissues remain in situ at 'ground-zero' where they are well positioned to conduct ongoing cancer immunosurveillance.

Previous studies have demonstrated in vitro that V\delta1 T cells have a Tc1 phenotype^{11,12,38,65}. However, given the scarcity of Vδ1

Fig. 3 | Intratumoral Vô1 T cells have a memory phenotype, are Tc1 skewed and demonstrate features of tissue residency and stemness. a,

Representative flow cytometry plots of effector memory status (defined by CD27 and CD45RA expression) of Vô1 T cells isolated from the NT tissue and tumor of one patient (representing n = 23 and n = 22 patients for NT tissue and tumor, respectively). **b**, Summary radar plot of effector memory status of Vô1 T cells isolated from NT tissues (n=23) and tumors (n=22). The median proportion is plotted. A two-tailed Mann-Whitney U-test was used between NT tissue and tumor within Vô1 memory subsets. Significant P values are shown. c, Expression of T cell master transcription factors and signature effector molecules of lymphocytes sorted directly from tumors grouped into T_H cell, cytolytic and inhibitory modules. Each column represents the denoted cell type from an individual patient. Not all cell types were sorted from matched patients. The color scale denotes the z-score of log₂(TPM + 1) of each gene. d, PCA of expression (normalized counts) of genes included in c colored by cell type (n=9, n=9, n=7, n=5, n=3 and n=5 patients for CD4⁺, CD8⁺, NK, T_{res}, Vδ2 and Vδ1 cells, respectively). **e**, Violin plots showing intracellular cytokine staining for IFN-γ and IL-17A and cell surface staining for CD107A in Vô1, CD8⁺ and CD4⁺ T cells after in vitro stimulation of bulk TILs (n = 3 patients) with PMA and ionomycin (P-I). f, Summary data of correlation between region-matched gene expression of NKG2D ligands and absolute numbers of V δ 1 (n=14 patients), V δ 2 (n=14 patients), CD4+ (n=15 patients) and CD8+ (n=15 patients) T_{RM} and T_{EM} cells in tumors. The color scale denotes a two-tailed Spearman's r. * denotes significant correlations as follows: MICA:V61 T_{EM} P=0.027, MICB:V&1 T_{EM} P=0.031, RAET1E:V&1 T_{EM} P=0.048 and RAET1E:CD8 T_{EM} P=0.045. **g**, PCA of expression (normalized counts) of core T_{RM} gene signature (CCR7, CD69, CXCR6, ITGAE, ITGA1, S1PR1 and SELL) in Vδ1 (n=5 patients), CD4+ (n=9 patients) and CD8+ T cells (n=9 patients) sorted from tumors. h, Expression of genes that define 'stem-like' CD8⁺ T cells in V δ 1 (n=5 patients), CD4⁺ (n=9 patients) and CD8⁺ (n=9 patients) T cells sorted from tumors. The mean ± s.d. is plotted. A Kruskal-Wallis test with post-hoc Dunn's test correction for multiple testing was used. All datapoints represent independent patients.

T cells, many of these studies have relied on ex vivo expansion using cytokine cocktails to achieve requisite numbers for in vitro assays. By using cells sorted directly from NSCLC tumors without ex vivo expansion or in vitro stimulation, we provide corroborating indirect and direct evidence for the in situ function of V δ 1 T cells through flow cytometric immunophenotyping and gene expression analysis. Compared with counterparts in NT lung tissues, V δ 1 T cells

isolated from tumors were enriched in the CD45RA⁻CD27⁻ T_{EM} subset that has previously been described as potent IFN- γ producers³⁸. Moreover, through gene expression analysis and in vitro functional assays, we demonstrate that intratumoral V δ 1 T cells are Tc1 skewed in situ.

Importantly, intratumoral V δ 1 T cells showed no evidence of a possibly tumor-promoting skew toward IL-17. Potentially



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Fig. 4 | Presence of Vd1 T cells associates with RFS in resected NSCLCs. a, RFS split on median absolute numbers of V δ 1, V δ 2, CD4⁺ and CD8⁺ T_{EM} cells in tumors. The Gehan-Breslow-Wilcoxon test was used. **b**, RFS split on median absolute numbers of V δ 1, V δ 2, CD4⁺ and CD8⁺ T_{EM} cells in NT tissues. The Gehan-Breslow-Wilcoxon test was used. **c**, Proportion of unique V δ 1 (*TRDV1*) T cell clones present in tumors and also found in paired NT tissues. The bar represents the median. A two-tailed Mann-Whitney *U*-test was used. **d**, Proportion of unique $\alpha\beta$ (TRA) T cell clones present in tumors and also found in paired NT tissues. The *n* numbers and datapoints represent independent patients.

dichotomous roles of $\gamma\delta$ T cells in cancer immunosurveillance and immunotherapy are currently contentious³⁷. Similar to $\alpha\beta$ T cells, $\gamma\delta$ T cells comprise distinct functional subsets. Thus, although absolute $\gamma\delta$ T cell deficiency predisposes to cancer in mice, it is now increasingly clear that T_H 1-cell-skewed, IFN- γ -producing $\gamma\delta$ T cells can be tumor rejecting⁵⁹ whereas T_H17-cell-skewed, IL-17-producing $\gamma\delta$ T cells may be tumor promoting^{18,66,67}. The limited evidence for IL-17 production by human $\gamma\delta$ T cells, as opposed to murine $\gamma\delta$ T cells, is confined mostly to a subset of peripheral blood-derived Vδ2 T cells⁴⁰⁻⁴². By contrast, human Vδ1 T cells have been consistently demonstrated to display a tumor-rejecting Tc1 phenotype when activated in vitro11,12,65 and now we provide evidence of this in situ within the TME. Consistent with this, we found a significant association of intratumoral T_{EM} V δ 1 T cells with ongoing remission. Although this association was based on outcomes in a modest cohort of patients with early stage, surgically resected NSCLCs, we found a similar association between high TRDV1 expression, as a proxy for intratumoral Vo1 T cells, and survival in public TCGA data of ~800 patients with NSCLCs. Moreover, the association of Vo1 T cells with favorable survival in TCGA was most evident in early-stage disease, alluding to the proposed role of $\gamma\delta$ T cells as proximal immune sentinels of epithelial stress⁵⁹. Our study does not directly address their role in metastatic disease where both the

larly at disparate metastatic sites, may be different. Likewise, we did not examine other immune compartments, such as the myeloid compartment or tertiary lymphoid structures, which may well contribute and warrant further study. Whether or not V δ 1 T cells directly effect tumor control or are merely a correlate of ongoing remission is difficult to answer without interventional studies such as adoptive V δ 1 T cell therapy, the first of which is currently under way for hematological cancers (NCT05001451). Nevertheless, the association of *TRDV1* expression with favorable responses to CPI therapy alludes to an important role for V δ 1 T cells. Indeed, a recent small study of mismatch repair-deficient colorectal cancers also implicated V δ 1 T cells in positive CPI responses⁶⁸.

tumor-intrinsic biology and immune microenvironment, particu-

In considering the utilization of the V δ 1 subset of $\gamma\delta$ T cells for cancer immunotherapy, one can now perceive several advantages over V δ 2 T cells. V δ 1 T cells are already resident in steady-state body surface tissues, raising the possibility that these cells may be beneficially manipulated in the clinic by off-the-shelf therapies, for example, via monoclonal antibodies targeting regulatory/stimulatory molecules⁶⁹. Within tumors themselves, V δ 1 T cells display a tissue-resident CD103⁺ phenotype, potentially improving their tumor-homing and tumor-retention capabilities vis-à-vis adoptive cell therapy. Finally, V δ 1 T cells have been demonstrated by others

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Fig. 5 | Expression of *TRDV1* **gene predicts NSCLC survival in TCGA and survival post-pembrolizumab in advanced solid cancers. a**, OS of patients with LUAD or LUSC in TCGA split on median *TRDV1* expression in primary tumor as a proxy for intratumoral V δ 1 T cells (n = 815 patients). The Gehan-Breslow-Wilcoxon test was used. **b**, OS of the same cohort of patients split on median *TRDC* expression as a proxy for intratumoral $\gamma\delta$ T cells (n = 815 patients). The Gehan-Breslow-Wilcoxon test was used. **b**, OS of the same cohort of patients split on median *TRDC* expression as a proxy for intratumoral $\gamma\delta$ T cells (n = 815 patients). The Gehan-Breslow-Wilcoxon test was used. NS, not significant. **c**, HRs for death in patients with above-median intratumoral expression of *TRDC* or *TRDV1* in UVA (n = 815 patients) and MVA (n = 774 patients) with age, gender, histology, smoking status, stage and *CD4* and *CD8B* gene expression. Rounded rectangles denote HRs and error bars denote 95% Cls. **d**, OS of patients with advanced solid cancers (mixed histologies) treated with pembrolizumab in the INSPIRE trial. There was a survival split on median *TRDV1* expression in primary tumor before pembrolizumab. The results were plotted in months after the third cycle of pembrolizumab. The Gehan-Breslow-Wilcoxon test was used. Significant *P* values are shown.

to be less susceptible to activation-induced cell death, a key barrier to durable responses in adoptive cell therapy⁷⁰.

Methods

Patients and samples. All clinical samples used were collected from patients recruited to the lung TRACERx Study (approved by an independent Research Ethics Committee, NRES Committee London, REC:13/LO/1546, https:// clinicaltrials.gov/ct2/show/NCT01888601). All participants provided informed consent before taking part. Participants were not compensated. Tissue specimens were reviewed by a lung pathologist as previously described²². Of note, NT tissues were taken as far away as possible from tumors at primary surgery and H&E sections examined afterwards by a trained histopathologist to ensure samples that were tumor free. Samples were chosen based on the availability of banked TILs from NT lung tissues and paired tumors, region-matched bulk DNA and patients with at least one follow-up visit after surgical resection. Where available, banked contemporaneous PBMCs were also immunophenotyped by flow cytometry. No other selection criteria were applied. Fresh NT tissue and NSCLCs were finely minced with sterile scalpels and dissociated in type 1 collagenase (10 U ml⁻¹, Thermo Fisher Scientific) and DNase I (75 µg ml-1, Roche) on a gentleMACS (Miltenyi Biotech) for 60 min at 37 °C. Digested material was passed through a 0.7-mm cell filter before TIL enrichment by Ficoll-paque gradient centrifugation (GE Healthcare). Isolated TILs were frozen in 10% dimethylsulfoxide (DMSO)/ fetal calf serum (FCS) and stored in liquid nitrogen until analysis. PBMCs were isolated from whole blood by Ficoll-paque gradient centrifugation, frozen in FCS with 10% DMSO (v:v) and stored in liquid nitrogen until analysis. DNA from paired tumor regions and paired NT tissues for TCR-seq was extracted as previously described²⁴. Briefly, region-matched tissues were homogenized using a TissueRuptor II (QIAGEN) and lysates passed through a QIAshredder column (QIAGEN) before DNA extraction using the Allprep DNA/RNA Mini kit (QIAGEN).

Flow cytometry and FACS. Thawed samples were washed in sterile phosphate-buffered saline (PBS) to remove traces of DMSO and serum before staining with Zombie NIR viability dye (1:500 dilution in PBS for 15 min at room temperature). Samples were then stained for lineage and differentiation markers for 15 min at 4°C (Supplementary Table 2; all antibodies used at 1:100 dilution in FACS buffer), washed twice with sterile 4°C FACS buffer, kept on ice and immediately acquired on a BD LSRFortessa or sorted on a BD FACSAria Fusion, running BD FACSDiva, and exported as FCS3.0 files. FCS3.0 files were analyzed on FlowJo v.10. For RNA-seq (see below), cells were sorted directly into ~4µl of lysis buffer (0.8% Triton X-100 in PBS (v:v) + 2 U ml⁻¹ of RNase inhibitor) at 4°C and lysates frozen at -80° C. Analysis and results were based on populations with a parent gate of ten or more cells.

In vitro activation assays. Thawed samples were washed in sterile PBS to remove traces of DMSO and rested overnight in complete RPMI medium (10% FCS + penicillin–streptomycin) at 37 °C and 5% CO₂. Rested cells were seeded at up to 200,000 cells per well in 200 µl of complete RPMI medium the next day. Cells either received no stimulation (complete RPMI medium only) or were stimulated with PMA (10 ng ml⁻¹) and ionomycin (1 µg ml⁻¹). Brefeldin A (5 µg ml⁻¹) and anti-CD107A (1:400 final dilution) were added to all the wells. Plates were centrifuged at 200g for 2 min and incubated at 37 °C and 5% CO₂

for 5 h. After 5 h, cells were stained for surface lineage markers as described above. After surface staining, samples were fixed in BD CellFIX and washed twice with permeabilization wash buffer (BioLegend) before staining for intracellular cytokines (1:100 final dilution of each antibody in permeabilization wash buffer) for 20 min at 4 °C. Samples were then washed twice with permeabilization wash buffer, resuspended in FACS buffer and immediately acquired on a BD LSRFortessa.

TCR-seq. The gDNA, 3 µg, extracted from region-matched tumors and NT tissues, was submitted for TCR α and TCR δ (TRA and TRD) sequencing with Adaptive Biotechnologies. Reads were aligned and annotated by Adaptive Biotechnologies. Sequences were filtered for in-frame CDR3 cells as well as TRA to TRA V-J family joins for $\alpha\beta$ T cells and TRD to TRD V-J family joins of $\gamma\delta$ T cells. Absolute counts of TCRs were normalized to 1 µg of input DNA for each sample to enable normalized comparison of T cell numbers across all samples. Data were analyzed using the immunoSEQ ANALYZER v.3.0 (Adaptive Biotechnologies).

RNA-seq of sorted TIL populations, data processing and PCAs. Where sufficient material was available, NK cells and T cell subsets (V81, V82, CD4+, CD8+ and T_{reg} cells) were sorted (50–500 cells from each subset) from NT tissues and tumors from a cohort of donors (patient IDs: CRUK0230, CRUK0299, CRUK0329, CRUK0344, CRUK0412, CRUK0416, CRUK0516, CRUK0860, CRUK0844, CRUK0926, CRUK0949, CRUK0968, CRUK0914, CRUK0961 and CRUK0922). Given the limited availability of material, not all cell types could be sorted from the same samples. Sorted cell lysates were frozen at -80 °C and submitted to the Oxford Genomics Centre for low-input library preparation and sequencing. Library preparation was completed from sorted cells in lysate buffer using the Smart-Seq2 protocol with minor modifications and NexteraXT (Illumina) following the manufacturer's instructions. Libraries were amplified (12 cycles) on a Tetrad (BioRad) using in-house dual indexing primers. Individual libraries were normalized using Qubit and the size profile was analyzed on the TapeStation 4200. Individual libraries were normalized and pooled together accordingly. The pooled library was diluted to ~10 nM for storage. The 10-nM library was denatured and further diluted before loading on the sequencer. Paired-end sequencing was performed using a HiSeq4000 75-bp platform (Illumina, HiSeq 3000/4000 PE Cluster Kit and 150-cycle SBS Kit), generating a raw read count of ~10 million reads per sample. The Trim Galore! utility v.0.4.2 (https://www.bioinformatics. babraham.ac.uk/projects/trim_galore-retrieved 3 May 2017) was used to remove sequencing adapters and quality trim individual reads with the q-parameter set to 20. Sequencing reads were aligned to the human genome and transcriptome (Ensembl GRCh38release-89) using RSEM v.1.3.0 (ref. 71) together with STAR aligner v.2.5.2 (ref. 72). Sequencing quality of individual samples was assessed using FASTQC v.0.11.5 (https://www.bioinformatics.babraham.ac.uk/projects/fastqcretrieved 3 May 17) and RNA-SeQC v.1.1.8 (ref. 73). PCAs were generated using normalized counts of the top 500 variable genes filtered for an average transcripts per million (TPM) across all samples >1 (Extended Data Fig. 3b) or normalized counts of curated master transcription factors and effector molecules presented in Fig. 3c (Fig. 3d).

Region-matched NKG2D ligand gene expression. For each region-matched sample, total RNA was extracted and prepared using a TruSeq Stranded Total RNA Human/Mouse/Rat ribo-depletion library preparation kit before Illumina complementary DNA paired-end sequencing. Libraries were prepared with

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 \geq 100 ng of total RNA input where possible and PCR amplified for 15 cycles. Libraries were quality checked by Agilent Tapestation and Promega QuantiFluor double-stranded DNA and pooled in equimolar amounts. Pooled libraries were sequenced in a HiSeq4000 at 50 million raw reads per sample, with a length of 75 bp or 100 bp per read. Illumina adapters were trimmed from raw reads using Cutadapt v.2.10 (ref. 74) with standard parameters. The quality of the trimmed reads was estimated per flow cell lane using FASTOC v.0.11.9. Fastq files with GC content within 2 s.d. of the cohort mean and <80% of total reads as duplicates were kept for alignment. These were then aligned to the UCSC hg19 human reference genome build using STAR aligner v.2.5.2a72 in two-pass mode with ENCODE 3 parameters generating one BAM file per tumor region. The same reads were also mapped to the human transcriptome (RefSeq GCA_000001405.1 build) using the same STAR parameters to generate gene expression data. Duplicates were marked with the MarkDuplicates function from GATK v.4.1.7.0 (ref. 75). Aligned reads were quality checked using QoRTs v.1.3.6 (ref. 76) for RNA integrity. Somalier v.0.2.7 (ref.) was used to detect potential instances of sample mislabeling. FASTQC, QoRTs and Somalier outputs were visualized using MultiQC v.1.9 (ref. 78). RSEM v.1.3.3 (ref. 71) was used with default parameters to quantify gene expression based on the BAM files aligned to the transcriptome. Gene expression patterns were used for further quality control of each sample. Tumor regions with <40% of all genes being expressed (>0 TPM) were excluded. In addition, samples with <20% of reads mapping uniquely to a single location in the genome were excluded.

TCGA analysis. NSCLC data from TCGA were acquired using R software (v.4.0.2). Gene expression data, Workflow Type: HTSeq-Counts and clinical data from the TCGA-LUAD and TCGA-LUSC projects were downloaded from Genomic Data Commons (GDC) Data Portal using the R/Bioconductor package TCGAbiolinks v.2.16.4 (ref. 79). A total of 594 lung adenocarcinoma (LUAD) and 551 lung squamous cell carcinoma (LUSC) cases were retrieved. Clinical data were utilized to select for primary tumor samples and for patients with OS of at least 1 d, yielding 474 LUAD and 473 LUSC cases. Samples in which TRDV1 transcript could be detected were included for analysis, thus yielding a final cohort of 417 LUAD and 398 LUSC cases (815 cases in total). Gene expression counts were normalized using DESeq2 v.1.28.1 (ref. 80) variance stabilizing transformation function and data were filtered for genes of interest: TRDV1 (V81 T cells) and TRDC (y8 cells). For survival analysis, patients were stratified into high- or low-expression groups for each gene of interest using median expression. OS was estimated from clinical data using 'days_to_last_followup' and 'days_to_death' and an OS event was defined from 'vital status' (Dead/Alive).

The association of *TRDV1* and *TRDC* with survival outcomes in NSCLCs was further assessed by multivariate analysis (MVA) adjusted for age (continuous), gender (categorical), histology (categorical), smoking status (categorical), stage (ordinal), *CD4* expression (above/below median) and *CD8B* expression (above/ below median) using the R package Survival (v.3.2-13). Of the variables assessed, only age, gender, histology, *CD4* expression and *CD8B* expression were available for all 815 cases. Smoking status was available for 782 cases and grouped into 'Ex-smokers', 'Current Smokers' and 'Never smokers' using 'tobacco_smoking_ history'. Patients with no smoking history information or current reformed smokers with no duration specified were excluded. Stage was available for 806 cases and grouped as stage 1, stage 2 and stage 3/4 owing to very few stage 4 cases. In total, 774 cases (out of 815 for *TRDV1/TRDC* univariate analysis (UVA)) were available for MVA.

GTEx analysis. The data used for the analyses described in Extended Data Fig. 2 were obtained from the GTEx portal (www.gtexportal.org/) and GTEx Analysis Release v.8 (dbGaP accession no. phs000424.v8,p2) based on a search for *TRDC/TRDV1/TRDV2* without any further selection criteria. The data were accessed on 14 January 2022. Detailed methods for the GTEx project are available from the GTEx portal.

Blood atlas analysis. The data used for analysis in Extended Data Fig. 5 were generated as part of the Blood Atlas Study³⁹. Gene expression data were downloaded from https://www.proteinatlas.org/about/download on 14 January 2022. TPM values were extracted from 'gdT cell', 'memory CD4 T cell', 'memory CD8 T cell', 'NK-cell' and 'T_{reg}' cell types, $\log_2(TPM + 1)$ transformed and z normalized across each gene.

INSPIRE survival analysis. Pre-processed RNA-seq and clinical data were downloaded for 51 patients in the INSPIRE cohort (NCT02644369) for whom both baseline gene expression and OS data were available. This pembrolizumab-treated cohort featured patients with a range of advanced solid cancers: head and neck (head and neck squamous cell cancers, n=9), TNBCs (n=6), high-grade serous carcinoma (n=8), melanoma (n=8) and other mixed solid tumors (n=20). TPM values that had been log₂(transformed) and batch normalized were downloaded as SourceData_Fig4.zip from Yang et al.⁵⁶ and survival split on median expression of each gene.

Statistics and reproducibility. No statistical method was used to predetermine sample size. Samples were chosen based on the availability of materials as described above. Three samples were excluded from outcome analysis due to involved

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The tumor region RNA-seq data, TCR-seq data and flow cytometry data (in each case from the TRACERx study) used or analyzed during the present study are available through the CRUK–University College London Cancer Trials Centre (ctc. tracerx@ucl.ac.uk) for academic noncommercial research purposes only, subject to review of a project proposal that will be evaluated by a TRACERx data access committee and any applicable ethical approvals, and entered into an appropriate data access agreement. Restrictions apply to the data availability to safeguard patient sequence data confidentiality, ensure compliance with patient study consent and meet data protection legislation, and due to commercial partnership requirements.

Details of all public datasets obtained from third parties used in the present study are as follows. Blood Atlas Study (https://doi.org/10.1126/science.aax9198) transcriptomic data were downloaded from https://www.proteinatlas.org/ about/download. GTEx (www.gtexportal.org) Analysis Release v.8 was accessed via dbGaP (accession no. phs000424.v8.p2). INSPIRE trial (NCT02644369) transcriptomic data were downloaded as SourceData_Fig4.zip from Yang et al. (https://doi.org/10.1038/s41467-021-25432-7). TCGA human LUAD and LUSC transcriptomic data were downloaded directly using the TCGAbiolinks R package derived from TCGA repository: https://portal.gdc.cancer.gov. Source data are provided with this paper.

Code availability

No customized code was used in the present study.

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Author contributions

Y.W. and D.B. designed and undertook laboratory experiments. Y.W., D.B., I.U., M.A., S.B., C.M., M.J., N.M. and A.H. designed, carried out and/or interpreted bioinformatics analyses and/or experimental data. T.K., S.V., J.C.K, S.H., J.R., A.G., M.A.-B., M.J.-H. and the TRACERx Consortium coordinated the clinical trial, and provided patient samples and patient data. Y.W., D.B., A.C.H. and C.S. wrote the manuscript. Y.W., S.A.Q., A.C.H and C.S. supervised the project. Y.W., A.C.H. and C.S. conceived and designed the project.

Competing interests

D.B. has consulted for NanoString, reports honoraria from AstraZeneca and has a patent (PCT/GB2020/050221) issued on methods for cancer prognostication. J.R. and M.A.B. have consulted for Achilles Therapeutics. N.M. has stock options in and has consulted for Achilles Therapeutics. N.M. holds European patents relating to targeting neoantigens (PCT/EP2016/059401), identifying patient response to immune checkpoint blockade (PCT/EP2016/071471), determining HLA loss of heterozygosity (PCT/GB2018/052004) and predicting survival rates of patients with cancer (PCT/GB2020/050221). A.H. attended one advisory board for Abbyle, Roche and GRAIL, and reports personal fees from Abbyle, Boehringer Ingelheim, Takeda, AstraZeneca, Daiichi Sankyo, Merck Serono, Merck/ MSD, UCB and Roche for delivering general education/training in clinical trials. A.H. owned shares in Illumina and Thermo Fisher Scientific (sold in 2020) and receives fees for membership of Independent Data Monitoring Committees for Roche-sponsored clinical trials. S.A.Q. is co-founder and Chief Scientific Officer of Achilles Therapeutics. A.C.H. is a board member and equity holder in ImmunoQure, AG and Gamma Delta Therapeutics, and is an equity holder in Adaptate Biotherapeutics and chair of the scientific advisory board. C.S. acknowledges grant support from Pfizer, AstraZeneca, Bristol Myers Squibb, Roche-Ventana, Boehringer Ingelheim, Archer Dx Inc (collaboration in minimal residual disease-sequencing technologies) and Ono Pharmaceuticals, is an AstraZeneca Advisory Board member and Chief Investigator for the MeRmaiD1 clinical trial. C.S has consulted for Amgen, AstraZeneca, Bicycle Therapeutics, Bristol Myers Squibb, Celgene, Genentech, GlaxoSmithKline, GRAIL, Illumina, Medixci, Metabomed, MSD, Novartis, Pfizer, Roche-Ventana and Sarah Cannon Research Institute. C.S. has stock options in Apogen Biotechnologies, Epic Biosciences and GRAIL, and has stock options and is co-founder of Achilles Therapeutics. C.S. holds patents relating: to assay technology to detect tumor recurrence (PCT/GB2017/053289); to targeting neoantigens (PCT/EP2016/059401), identifying patent response to immune checkpoint blockade (PCT/EP2016/071471), determining HLA loss of heterozygosity (PCT/GB2018/052004), predicting survival rates of patients with cancer (PCT/GB2020/050221); to treating cancer by targeting Insertion/ deletion (indel) mutations (PCT/GB2018/051893); to identifying indel mutation targets (PCT/GB2018/051892); to methods for lung cancer detection (PCT/US2017/028013); and to identifying responders to cancer treatment (PCT/GB2018/051912). The remaining authors declare no competing interests.

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Correspondence and requests for materials should be addressed to Yin Wu, Adrian C. Hayday or Charles Swanton.

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TRACERx Consortium

Charles Swanton^{1,2}, Mariam Jamal-Hanjani^{1,7}, Nicholas McGranahan^{1,9}, Allan Hackshaw¹⁰, Yin Wu^{1,2,3,4,55}, Dhruva Biswas^{1,2,5,55}, Mihaela Angelova², Stefan Boeing⁶, Takahiro Karasaki^{1,2}, Selvaraju Veeriah¹, Sonya Hessey^{1,7}, James Reading^{1,8}, Maise Al-Bakir², Sergio A. Quezada^{1,8}, Nicolai J. Birkbak¹¹, Gillian Price¹², Mohammed Khalil¹², Keith Kerr¹², Shirley Richardson¹², Heather Cheyne¹², Tracey Cruickshank¹², Gareth A. Wilson¹³, Rachel Rosenthal¹³, Hugo Aerts^{14,15}, Madeleine Hewish¹⁶, Girija Anand¹⁷, Sajid Khan¹⁷, Kelvin Lau¹⁸, Michael Sheaff¹⁸, Peter Schmid¹⁸, Louise Lim¹⁸, John Conibear¹⁸, Roland Schwarz¹⁹, Tom L. Kaufmann¹⁹, Matthew Huska¹⁹, Jacqui Shaw²⁰,

Joan Riley²⁰, Lindsay Primrose²⁰, Dean Fennell^{20,21}, Allan Hackshaw²², Yenting Ngai²², Abigail Sharp²², Oliver Pressey²², Sean Smith²², Nicole Gower²², Harjot Kaur Dhanda²², Kitty Chan²², Sonal Chakraborty²², Kevin Litchfield¹, Krupa Thakkar¹, Jonathan Tugwood²³, Alexandra Clipson²³, Caroline Dive^{23,24}, Dominic Rothwell^{23,24}, Alastair Kerr^{23,24}, Elaine Kilgour^{23,24}, Fiona Morgan²⁵, Malgorzata Kornaszewska²⁵, Richard Attanoos²⁵, Helen Davies²⁵, Katie Baker²⁶, Mathew Carter²⁶, Colin R. Lindsay²⁶, Fabio Gomes²⁶, Fiona Blackhall^{24,26}, Lynsey Priest^{24,26}, Matthew G. Krebs^{24,26}, Anshuman Chaturvedi^{24,26}, Pedro Oliveira^{24,26}, Zoltan Szallasi²⁷, Gary Royle²⁸, Catarina Veiga²⁸, Marcin Skrzypski²⁹, Roberto Salgado³⁰, Miklos Diossy³¹, Alan Kirk³², Mo Asif³², John Butler³², Rocco Bilancia³², Nikos Kostoulas³², Mathew Thomas³², Mairead MacKenzie³³, Maggie Wilcox³³, Apostolos Nakas²¹, Sridhar Rathinam²¹, Rebecca Boyles²¹, Mohamad Tufail²¹, Amrita Bajaj²¹, Keng Ang²¹, Mohammed Fiyaz Chowdhry²¹, Michael Shackcloth³⁴, Julius Asante-Siaw³⁴, Angela Leek³⁵, Nicola Totten³⁵, Jack Davies Hodgkinson³⁵, Peter Van Loo³⁶, William Monteiro³⁷, Hilary Marshal³⁷, Kevin G. Blyth³⁸, Craig Dick³⁸, Charles Fekete¹⁵, Eric Lim³⁹, Paulo De Sousa³⁹, Simon Jordan³⁹, Alexandra Rice³⁹, Hilgardt Raubenheimer³⁹, Harshil Bhayani³⁹, Morag Hamilton³⁹, Lyn Ambrose³⁹, Anand Devaraj³⁹, Hemangi Chavan³⁹, Sofina Begum³⁹, Silviu I. Buderi³⁹, Daniel Kaniu³⁹, Mpho Malima³⁹, Sarah Booth³⁹, Andrew G. Nicholson³⁹, Nadia Fernandes³⁹, Pratibha Shah³⁹, Chiara Proli³⁹, John Gosney⁴⁰, Sarah Danson⁴¹, Jonathan Bury⁴¹, John Edwards⁴¹, Jennifer Hill⁴¹, Sue Matthews⁴¹, Yota Kitsanta⁴¹, Jagan Rao⁴¹, Sara Tenconi⁴¹, Laura Socci⁴¹, Kim Suvarna⁴¹, Faith Kibutu⁴¹, Patricia Fisher⁴¹, Robin Young⁴¹, Joann Barker⁴¹, Fiona Taylor⁴¹, Kirsty Lloyd⁴¹, Jason Lester⁴², Mickael Escudero⁴³, Aengus Stewart⁴³, Andrew Rowan⁴³, Jacki Goldman⁴³, Richard Kevin Stone⁴³, Tamara Denner⁴³, Emma Nye⁴³, Maria Greco⁴³, Jerome Nicod⁴³, Clare Puttick⁴³, Katey Enfield⁴³, Emma Colliver⁴³, Alastair Magness⁴³, Chris Bailey⁴³, Krijn Dijkstra⁴³, Vittorio Barbè⁴³, Roberto Vendramin⁴³, Judit Kisistok⁴³, Mateo Sokac⁴³, Jonas Demeulemeester⁴³, Elizabeth Larose Cadieux⁴³, Carla Castignani⁴³, Hongchang Fu⁴³, Kristiana Grigoriadis⁴³, Claudia Lee^{43,44}, Foteini Athanasopoulou^{1,43}, Crispin Hiley^{1,43}, Lily Robinson⁴⁵, Tracey Horey⁴⁵, Peter Russell⁴⁵, Dionysis Papadatos-Pastos⁴⁵, Sara Lock⁴⁶, Kayleigh Gilbert⁴⁶, Kayalvizhi Selvaraju⁴⁷, Paul Ashford⁴⁷, Oriol Pich⁴⁷, Thomas B. K. Watkins⁴⁷, Sophia Ward⁴⁷, Emilia Lim⁴⁷, Alexander M. Frankell⁴⁷, Christopher Abbosh⁴⁷, Robert E. Hynds⁴⁷, Mariana Werner Sunderland⁴⁷, Karl Peggs⁴⁷, Teresa Marafioti⁴⁷, John A. Hartley⁴⁷, Helen Lowe⁴⁷, Leah Ensell⁴⁷, Victoria Spanswick⁴⁷, Angeliki Karamani⁴⁷, David Moore⁴⁷, Stephan Beck⁴⁷, Olga Chervova⁴⁷, Miljana Tanic⁴⁷, Ariana Huebner⁴⁷, Michelle Dietzen⁴⁷, James R. M. Black⁴⁷, Carlos Martinez Ruiz⁴⁷, Robert Bentham⁴⁷, Cristina Naceur-Lombardelli⁴⁷, Haoran Zhai⁴⁷, Nnennaya Kanu⁴⁷, Francisco Gimeno-Valiente⁴⁷, Supreet Kaur Bola⁴⁷, Ignacio Garcia Matos⁴⁷, Mansi Shah⁴⁷, Felipe Galvez Cancino⁴⁷, Despoina Karagianni⁴⁷, Maryam Razaq⁴⁷, Mita Akther⁴⁷, Diana Johnson⁴⁷, Joanne Laycock⁴⁷, Elena Hoxha⁴⁷, Benny Chain⁴⁷, David R. Pearce⁴⁷, Kezhong Chen⁴⁷, Javier Herrero⁴⁷, Fleur Monk⁴⁷, Simone Zaccaria⁴⁷, Neil Magno⁴⁷, Paulina Prymas⁴⁷, Antonia Toncheva⁴⁷, Monica Sivakumar⁴⁷, Olivia Lucas^{47,48}, Mark S. Hill^{43,47}, Othman Al-Sawaf⁴⁷, Seng Kuong Ung⁴⁸, Sam Gamble⁴⁸, Sophia Wong⁴⁸, David Lawrence⁴⁸, Martin Hayward⁴⁸, Nikolaos Panagiotopoulos⁴⁸, Robert George⁴⁸, Davide Patrini⁴⁸, Mary Falzon⁴⁸, Elaine Borg⁴⁸, Reena Khiroya⁴⁸, Asia Ahmed⁴⁸, Magali Taylor⁴⁸, Junaid Choudhary⁴⁸, Sam M. Janes⁴⁸, Martin Forster⁴⁸, Tanya Ahmad⁴⁸, Siow Ming Lee⁴⁸, Neal Navani⁴⁸, Marco Scarci⁴⁸, Pat Gorman⁴⁸, Elisa Bertoja⁴⁸, Robert C. M. Stephens⁴⁸, Emilie Martinoni Hoogenboom⁴⁸, James W. Holding⁴⁸, Steve Bandula⁴⁸, Ricky Thakrar⁴⁸, James Wilson⁴⁸, Mansi Shah⁴⁸, Marcos⁴⁸, Vasquez Duran⁴⁸, Maria Litovchenko⁴⁸, Sharon Vanloo⁴⁸,

Piotr Pawlik⁴⁸, Kerstin Thol⁴⁸, Babu Naidu⁴⁹, Gerald Langman⁴⁹, Hollie Bancroft⁴⁹, Salma Kadiri⁴⁹, Gary Middleton⁴⁹, Madava Djearaman⁴⁹, Aya Osman⁴⁹, Helen Shackleford⁴⁹, Akshay Patel⁴⁹, Christian Ottensmeier⁵⁰, Serena Chee⁵⁰, Aiman Alzetani⁵⁰, Judith Cave⁵⁰, Lydia Scarlett⁵⁰, Jennifer Richards⁵⁰, Papawadee Ingram⁵⁰, Emily Shaw⁵⁰, John Le Quesne⁵¹, Alan Dawson⁵², Domenic Marrone⁵², Sean Dulloo⁵², Claire Wilson⁵², Yvonne Summers⁵³, Raffaele Califano⁵³, Rajesh Shah⁵³, Piotr Krysiak⁵³, Kendadai Rammohan⁵³, Eustace Fontaine⁵³, Richard Booton⁵³, Matthew Evison⁵³, Stuart Moss⁵³, Juliette Novasio⁵³, Leena Joseph⁵³, Paul Bishop⁵³, Helen Doran⁵³, Felice Granato⁵³, Vijay Joshi⁵³, Elaine Smith⁵³, Angeles Montero⁵³ and Phil Crosbie^{53,54}

¹¹Aarhus University, Aarhus, Denmark. ¹²Aberdeen Royal Infirmary, Aberdeen, UK. ¹³Achilles Therapeutics UK Limited, London, UK. ¹⁴Artificial Intelligence in Medicine Program, Harvard Medical School, Boston, MA, USA. ¹⁵Radiology and Nuclear Medicine, CARIM & GROW, Maastricht University, Maastricht, The Netherlands. ¹⁶Ashford and St. Peter's Hospitals NHS Foundation Trust, Chertsey, UK. ¹⁷Barnet & Chase Farm Hospitals, London, UK. ¹⁸Barts Health NHS Trust, London, UK. ¹⁹Berlin Institute for Medical Systems Biology, Max Delbrück Center for Molecular Medicine in the Helmholtz Association, Berlin, Germany. ²⁰Cancer Research Centre, University of Leicester, Leicester, UK. ²¹Leicester University Hospitals, Leicester, UK. ²²Cancer Research UK & UCL Cancer Trials Centre, London, UK. ²³Cancer Research UK Manchester Institute, University of Manchester, Manchester, UK. ²⁴Cancer Research UK Lung Cancer Centre of Excellence, University of Manchester, Manchester, UK. 25 Cardiff & Vale University Health Board, Cardiff, UK. 26 Christie NHS Foundation Trust, Manchester, UK. ²⁷Danish Cancer Society Research Centre, Copenhagen, Denmark. ²⁸Department of Medical Physics and Bioengineering, University College London Cancer Institute, London, UK. ²⁹Department of Oncology and Radiotherapy, Medical University of Gdańsk, Gdańsk, Poland. ³⁰Department of Pathology, GZA-ZNA Antwerp, Antwerp, Belgium. ³¹Department of Physics of Complex Systems, ELTE Eötvös Loránd University, Budapest, Hungary. ³²Golden Jubilee National Hospital, Clydebank, UK. ³³Independent Cancer Patients Voice, London, UK. ³⁴Liverpool Heart and Chest Hospital NHS Foundation Trust, Liverpool, UK.³⁵Manchester Cancer Research Centre Biobank, Manchester, UK.³⁶MD Anderson Cancer Center, Houston, TX, USA. ³⁷National Institute for Health Research, Leicester Respiratory Biomedical Research Unit, Leicester, UK. ³⁸NHS Greater Glasgow and Clyde, Glasgow, UK. ³⁹Royal Brompton and Harefield NHS Foundation Trust, London, UK. ⁴⁰Royal Liverpool University Hospital, Liverpool, UK. ⁴¹Sheffield Teaching Hospitals NHS Foundation Trust, Sheffield, UK. ⁴²Swansea Bay University Health Board, Swansea, UK. ⁴³The Francis Crick Institute, London, UK. ⁴⁴University College London Medical School, London, UK. ⁴⁵The Princess Alexandra Hospital NHS Trust, Harlow, UK. ⁴⁶The Whittington Hospital NHS Trust, London, UK. ⁴⁷University College London Cancer Institute, London, UK. ⁴⁸University College London Hospitals, London, UK. ⁴⁹University Hospital Birmingham NHS Foundation Trust, Birmingham, UK. 50 University Hospital Southampton NHS Foundation Trust, Southampton, UK. 51 Institute of Cancer Sciences, University of Glasgow, Glasgow, UK. 52 University of Leicester, Leicester, UK. 53 Wythenshawe Hospital, Manchester University NHS Foundation Trust, Manchester, UK. ⁵⁴Division of Infection, Immunity and Respiratory Medicine, University of Manchester, Manchester, UK.

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Extended Data Fig. 1 | Lung $\gamma\delta$ **T cells assayed by flow cytometry and TCR sequencing. A)** Representative flow cytometry gating strategy for T cell subsets. From left to right: Forward vs. side scatter gate -> singlet gate -> live CD45 + gate -> CD3 vs. TCR $\gamma\delta$ -> CD4 vs. CD8 (of CD3 +/TCR $\gamma\delta$ -population) and V $\delta1$ vs. V $\delta2$ (of CD3 +/TCR $\gamma\delta$ + population) to give CD4 and CD8 $\alpha\beta$ T cells and V $\delta1$ and V $\delta2$ $\gamma\delta$ T cells. **B)** Representative gating strategy for CD103 and CD45RA vs. CD27 for V $\delta1$ and V $\delta2$ $\gamma\delta$ T cells, and CD4 and CD8 $\alpha\beta$ T cells. **C)** Proportion of T cell subsets determined by flow cytometry versus proportion determined by TCRseq in NT tissues (n = 23 patients) and tumours (n = 22 patients). Two-tailed Spearman correlation. Significant p values shown. All datapoints represent independent patients.

ARTICLES



Extended Data Fig. 2 | TCR δ is expressed in non-diseased lung tissues. A) Expression of *TRDC* across non-diseased tissue types. B) Expression of *TRDV1* across non-diseased tissue types. C) Expression of *TRDV2* across non-diseased tissue types. Gene expression data and figures derived from the GTEx Project and Portal (see methods). Lung expression (red arrow) based on 578 donors. Boxplots represent median and interquartile range with outliers (above or below 1.5x IQR) displayed as points. Tissues ordered from highest to lowest median expression of each gene. Datapoints represent independent donors.

NATURE CANCER



Extended Data Fig. 3 | Lung TIL lineages by RNA sequencing. A) Gating strategy for sorting CD4, CD8, V δ 1 and V δ 2 T cells and Tregs and NK cells (red/bold) for bulk cell-type RNA sequencing. B) PCA of top 500 variable genes coloured by cell type from tumours (n = 9, n = 9, n = 7, n = 5, n = 3 and n = 5 patients for CD4⁺, CD8⁺, NK, Treg, V δ 2 and V δ 1 cells respectively) and NT tissue (n = 8, n = 7, n = 8, n = 1, n = 2 and n = 2 patients for CD4⁺, CD8⁺, NK, Treg, V δ 2 and V δ 1 cells respectively) and NT tissue (n = 8, n = 7, n = 8, n = 1, n = 2 and n = 2 patients for CD4⁺, CD8⁺, NK, Treg, V δ 2 and V δ 1 cells respectively). C) Expression of canonical lineage markers from sorted cell types. Lineage markers for each sorted cell type coloured in. Cells sorted from NT tissues denoted by crossed-circles (n = 8, n = 7, n = 8, n = 1, n = 2 and n = 2 patients for CD4⁺, CD8⁺, NK, Treg, V δ 2 and V δ 1 cells respectively) and cells sorted from lung tumours denoted by solid circles (n = 9, n = 9, n = 7, n = 5, n = 3 and n = 5 patients for CD4⁺, CD8⁺, NK, Treg, V δ 2 and V δ 1 cells respectively). All datapoints represent independent patients.



Extended Data Fig. 4 | Peripheral blood γδ **T cell functional transcriptomes.** Expression of T cell master transcription factors and signature effector molecules of peripheral blood lymphocytes from the Blood Atlas project. Each column represents the denoted cell type from an individual donor. Colour scale denotes Z-score of Log2 (TPM +1) of each gene.

NATURE CANCER



Extended Data Fig. 5 | Effector function of *in vitro* stimulated TILs from NSCLCs. Representative flow cytometry dot plots of intracellular cytokine staining for IFN- γ and IL-17A and cell surface staining for CD107A in V δ 1, CD8 and CD4 T cells after *in vitro* stimulation of bulk tumour infiltrating lymphocytes with PMA and ionomycin (P-I). Gates were set on paired unstimulated negative controls (-). Percentage positive of parent population shown.

ARTICLES



Extended Data Fig. 6 | Tumour Vδ1 T cells demonstrate no evidence of clonotypic responses. A) Repertoire analysis by normalised Shannon entropy (S.E.) of Vδ1 T cells in NT tissue (Tis) and paired lung tumours (Tum) calculated on TCRδ CDR3 amino-acid sequences. Wilcoxon matched-pairs signed rank test. **B)** Repertoire analysis by D50 of Vδ1 T cells in NT tissue (Tis) and paired lung tumours (Tum) calculated on TCRδ CDR3 amino-acid sequences. Wilcoxon matched-pairs signed rank test. **C)** Relative expression (TPM) of functional TCRγ Vγ genes in sorted intra- tumoural Vδ1 T cells from 5 patients and Vδ2 from 3 patients. All datapoints represent independent patients.

NATURE CANCER



Extended Data Fig. 7 | Tumour Vô1 T cells possess a core T_{RM} **signature.** Expression of core TRM signature genes in Vô1, CD8, and CD4 T cells sorted from tumours (n = 5, n = 9 and n = 9 patients respectively). Genes associated with tissue retention and homing and expected to be upregulated in TRM T cells highlighted in red. Genes associated with tissue egress and expected to be downregulated in TRM T cells highlighted in blue. Mean +/- S.D. plotted. Kruskal Wallis with post hoc Dunn's test corrected for multiple testing. All datapoints represent independent patients.

ARTICLES



Extended Data Fig. 8 | Prognostic value of Vô1 T cells is largely independent of other prognostic clinicopathological features. A) Absolute numbers of Vô1 TEM cells in tumours plotted by TNM stage, histology and smoking status (never smoker, unfilled circle) as well as against age and primary size. Kruskal Wallis with post hoc Dunn's test corrected for multiple testing (stage). Bar=median. Mann-Whitney test (histology and smoking status). Spearman correlation (primary size and age). B) Relapse free survival split on median absolute numbers of Vô1 Trm cells in tumours. Gehan-Breslow-Wilcoxon test. **C)** Absolute numbers of Vô1 TEM cells in NT tissues plotted by TNM stage, histology and smoking status (never smoker, unfilled circle) as well as against age and primary size. Bar=median. Kruskal Wallis with post hoc Dunn's test corrected for multiple testing (stage). Mann-Whitney test (histology and smoking status). Spearman correlation (primary size and age). **D)** Δ Vô1 TEM cells (absolute Vd1 TEM cells in tumour - absolute Vd1 TEM cells in paired NT tissue) and Δ Vô1 TRM cells (absolute Vd1 TRM cells in tumour - absolute Vd1 TRM cells in paired NT plotted by outcome. Bar=median. Mann-Whitney test. N numbers and datapoints represent independent patients.

LES **NATURE CANCER** INSPIRE TRDC **INSPIRE** TRDV2 **INSPIRE** CD4 **INSPIRE** CD8B 100 100 100 100 p=0.097 p=0.55 p=0.062 80 p=0.025 80 80 80 Percent OS Percent OS n.s. Percent OS n.s. Percent OS n.s 60 60 60 60 40 40 40 40 – Low (n=25) – High (n=26) Low (n=25) High (n=26) – Low (n=25) – High (n=26) 20 20 20 20 Low (n=25) High (n=26) 0↓ 0 0↓ 0 0↓ 0 0↓ 0 10 15 20 25 30 35 10 15 20 25 30 35 10 15 20 25 30 35 10 15 20 25 30 35 5 5 5 5 Months Months Months Months

Extended Data Fig. 9 | Association of survival with signature T cell lineage genes in the INSPIRE Trial. Overall survival of patients with advanced solid cancers (mixed histologies) treated with pembrolizumab in the INSPIRE trial. Survival split on median TRDC, TRDV2, CD4 and CD8B expression in primary tumour prior to pembrolizumab. Plotted in months after third cycle of pembrolizumab. Gehan-Breslow-Wilcoxon test. Significant p values shown. n.s. not significant.

nature portfolio

Yin Wu Adrian Hayday Corresponding author(s): Charles Swanton

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Reporting Summary

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Statistics

For	all st	atistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.
n/a	Cor	firmed
	\boxtimes	The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
\boxtimes		A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
	\boxtimes	The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.
	\boxtimes	A description of all covariates tested
	\boxtimes	A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
	\boxtimes	A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
		For null hypothesis testing, the test statistic (e.g. <i>F</i> , <i>t</i> , <i>r</i>) with confidence intervals, effect sizes, degrees of freedom and <i>P</i> value noted <i>Give P values as exact values whenever suitable.</i>
\boxtimes		For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
\boxtimes		For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
	\boxtimes	Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated
		Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.

Software and code

Policy information about <u>availability of computer code</u>		
Data collection	No custom code was used to collect data.	
	NSCLC data from TCGA was acquired using R software (version 4.0.2). Gene expression data, Workflow Type: HTSeq-Counts and clinical data from the TCGA-LUAD and TCGA-LUSC projects were downloaded from Genomic Data Commons (GDC) Data Portal using the R/Bioconductor package TCGAbiolinks version 2.16.4.	
	BD FACSDiva software was used for collection of flow cytometry data	
Data analysis	No custom code was used for data analysis.	
	CCutadapt version 2.10	
	DESeq2 version 1.28.185	
	FASTQC version 0.11.5	
	FASTQC version 0.11.9	
	GATK version 4.1.7.0	
	MultiQC version 1.9	
	QoRTs version 1.3.6	
	RNA-SeQC version 1.1.8	
	RSEM version 1.3.0	
	RSEM version 1.3.3	
	Somalier version 0.2.7	
	STAR aligner version 2.5.2	
	STAR aligner version 2.5.2a	
	Survival version 3.2-13	

Trim Galore! utility version 0.4.2 FlowJo version 10 GraphPad Prism version 9.2.0 immunoSEQ Analyzer version 3.0 JMP Pro version 15.0.0

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio guidelines for submitting code & software for further information.

Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A description of any restrictions on data availability
- For clinical datasets or third party data, please ensure that the statement adheres to our policy

The tumour region RNA-seq data, TCR-seq data, and flow cytometry data (in each case from the TRACERx study), used or analysed during this study are available through the Cancer Research UK & University College London Cancer Trials Centre (ctc.tracerx@ucl.ac.uk) for academic non-commercial research purposes only, subject to review of a project proposal that will be evaluated by a TRACERx data access committee and any applicable ethical approvals, and entering into an appropriate data access agreement. Restrictions apply to the data availability in order to safeguard patient sequence data confidentiality, to ensure compliance with patient study consent, to meet data protection legislation and due to commercial partnership requirements.

Details of all public datasets obtained from third parties used in this study are as follows. Blood Atlas Study (Uhlen et al. 2019, doi.org/10.1126/science.aax9198) transcriptomic data was downloaded from https://www.proteinatlas.org/about/download. GTEx (www.gtexportal.org/) Analysis Release V8 was accessed via dbGaP Accession phs000424.v8.p2. INSPIRE trial (NCT02644369) transcriptomic data were downloaded as SourceData_Fig4.zip from Yang et al. 2021, doi.org/10.1038/ s41467-021-25432-7. TCGA human LUAD and LUSC transcriptomic data were downloaded directly using the TCGAbiolinks R package derived from TCGA repository https://portal.gdc.cancer.gov/.

Source data are available as Source Data files.

Field-specific reporting

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Life sciences

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Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see <u>nature.com/documents/nr-reporting-summary-flat.pdf</u>

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	Samples (n=25) were chosen based on the availability of banked TILs from NT lung tissue and paired tumours, region-matched bulk DNA, and from patients with at least one follow-up visit after surgical resection. Where available, banked contemporaneous PBMCs were also immunophenotyped by flow cytometry. No other selection criteria were applied.
Data exclusions	Three patients were excluded from outcome analysis due to incomplete surgical resection of their primary tumours.
Replication	Survival outcome was validated in a cohort of 800+ patients with non-small cell lung cancer in the TCGA database. Technical replicates for flow cytometry, TCR sequencing, RNA sequencing and in vitro activation assays were not performed due to the limited availability of material. Biological replicates from multiple independent donors cross-validated the presence of gamma-delta T cell subsets and their phenotypes in clinical samples.
Randomization	Non-interventional study, not applicable
Blinding	Non-interventional study, not applicable

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

Methods

ChIP-seq

n/a Involved in the study

Flow cytometry

MRI-based neuroimaging

n/a	Involved in the study
	X Antibodies
\boxtimes	Eukaryotic cell lines
\boxtimes	Palaeontology and archaeology
\boxtimes	Animals and other organisms
	Human research participants
	🔀 Clinical data
\boxtimes	Dual use research of concern

Antibodies

Antibodies used	Anti-CD3 (PerCP-CyS.5) Biolegend Cat# 317336 Dilution 1:100 Anti-CD4 (PE-Dazzle594) Biolegend Cat# 317448 Dilution 1:100 Anti-CD8a (AF700) Biolegend Cat# 30920 Dilution 1:100 Anti-CD14 (PE-Cy7) Biolegend Cat# 367112 Dilution 1:100 Anti-CD19 (AF700) Biolegend Cat# 363033 Dilution 1:100 Anti-CD25 (BV650) Biolegend Cat# 302634 Dilution 1:100 Anti-CD27 (BV786) BD Biosciences Cat# 563327 Dilution 1:100 Anti-CD45 (BV605) Biolegend Cat# 304042 Dilution 1:100 Anti-CD45 (BV605) Biolegend Cat# 304042 Dilution 1:100 Anti-CD56 (APC) Biolegend Cat# 302144 Dilution 1:100 Anti-CD56 (APC) Biolegend Cat# 350214 Dilution 1:100 Anti-CD103 (BV421) Biolegend Cat# 350214 Dilution 1:100 Anti-CD107A (BV650) Biolegend 512333 Dilution 1:100 Anti-L1-T7 (APC) Biolegend 502531 Dilution 1:100 Anti-TCR Vδ1 (FITC) ThermoFisher Cat# TCR2730 Dilution 1:100 Anti-TCR Vδ2 (PE) Biolegend 3Cat# 31048 Dilution 1:100
Validation	Antibodies and reagents used have been validated by commercial suppliers and validation statements can be found on suppliers' websites. Biolegend https://www.biolegend.com/en-us/quality/quality-control "All of our products undergo industry-leading rigorous quality control (QC) testing to ensure the highest level of performance and reproducible results. Each lot is compared to an internally established "gold standard" to maintain lot-to-lot consistency. We also conduct wide-scale stability studies to guarantee an accurate shelf-life for our products. Additionally, we test the majority of our products on endogenous cells rather than transfected or immortal cells that may overexpress the analyte. We assess our reagents with samples and protocols that reflect our customers' experience. Our willingness to monitor the quality of our reagents extends beyond our lab and into yours."
	BD Biosciences https://www.bdbiosciences.com/en-gb/products/reagents/flow-cytometry-reagents/research-reagents/quality-and-reproducibility "BD Biosciences not only develops its own antibodies but also collaborates with research scientists around the world to license their antibodies. We provide accessibility to the flow cytometry community by conjugating antibodies to a broad portfolio of high- performing dyes, including our vastly popular portfolio of BD Horizon BrilliantTM Dyes. A world-class team of research scientists helps ensure that these reagents work reliably and consistently for flow cytometry applications. The specificity is confirmed using multiple applications that may include a combination of flow cytometry, immunofluorescence, immunohistochemistry or western blot to test a combination of primary cells, cell lines or transfectant models. All flow cytometry reagents are titrated on the relevant positive or negative cells. To save time and cell samples for researchers, pre-titrated test size reagents are bottled at an optimal concentration, with the best signal-to-noise ratio on relevant models."
	https://www.mybeckman.uk/reagents/coulter-flow-cytometry "Our portfolio of Flow Cytometry reagents, entirely manufactured under good manufacturing practices (GMP), covers major application areas including hemato-oncology, HIV analysis, immune monitoring, cell cycle and stem cells studies."
	ThermoFisher https://www.thermofisher.com/uk/en/home/life-science/antibodies/invitrogen-antibody-validation.html "Part 1—Target specificity verification This helps ensure the antibody will bind to the correct target. Our antibodies are being tested using at least one of the following methods to ensure proper functionality in researcher's experiments. Click on each testing method below for detailed testing strategies, workflow examples, and data figure legends.
	Knockout—expression testing using CRISPR-Cas9 cell models Knockdown—expression testing using RNAi to knockdown gene of interest Independent antibody verification (IAV)—measurement of target expression is performed using two differentially raised antibodies

recognizing the same protein target
Cell treatment—detecting downstream events following cell treatment
Relative expression—using naturally occurring variable expression to confirm specificity
Neutralization—functional blocking of protein activity by antibody binding
Peptide array—using arrays to test reactivity against known protein modifications
SNAP-ChIP™—using SNAP-ChIP to test reactivity against known protein modifications
Immunoprecipitation-Mass Spectrometry (IP-MS)—testing using immunoprecipitation followed by mass spectrometry to identify
antibody targets
Part 2—Functional application validation
These tests help ensure the antibody works in a particular application(s) of interest, which may include
(but are not limited to):
Western blotting
Flow cytometry
ChIP
Immunofluorescence imaging
Immunohistochemistry
Most antibodies were developed with specific applications in mind. Testing that an antibody generates acceptable results in a specific
application is the second part of confirming antibody performance."

Human research participants

Policy information about studies involving human research participants

Population characteristics	This study includes a subset of patients recruited to the TRACERx Study for which population characteristics have previously been reported (DOI: 10.1056/NEJMoa1616288).
	The population characteristics for the subset of patients included in this study are detailed in Supplementary Table 1
Recruitment	Samples (n=25) for this study were chosen based on the availability of banked TILs from NT lung tissue and paired tumours, region-matched bulk DNA, and from patients with at least one follow-up visit after surgical resection. Where available, banked contemporaneous PBMCs were also immunophenotyped by flow cytometry. No other selection criteria were applied. These criteria may potentially select for larger primary tumours within each stage and results should be interpreted with this in mind.
Ethics oversight	The study was approved by the NRES Committee London - Camden and Islington with the following details: Study title: TRAcking non small cell lung Cancer Evolution through therapy (Rx) REC reference: 13/LO/1546 Protocol number: UCL/12/0279 IRAS project ID: 138871
	All participants provided informed consent prior to taking part.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

Clinical data

Policy information about <u>clinical studies</u> All manuscripts should comply with the ICMJE <u>guidelines for publication of clinical research</u> and a completed <u>CONSORT checklist</u> must be included with all submissions.

Clinical trial registration	TRACERx Lung: https://clinicaltrials.gov/ct2/show/NCT01888601, approved by an independent Research Ethics Committee, 13/ LO/1546 INSPIRE Trial: https://clinicaltrials.gov/ct2/show/NCT02644369
Study protocol	TRACERx Lung: The study protocol is available as a supplement to Jamal-Hanjani et al. 2017, https://doi.org/10.1056/ NEJMoa1616288
	INSPIRE Trial: The study protocol is as a supplement to Yang et al. 2021, https://doi.org/10.1038/s41467-021-25432
Data collection	TRACERx Lung: Clinical and pathological data is collected from patients during study follow up - this period is a minimum of five years. Data collection is overseen by the sponsor of the study (Cancer Research UK & UCL Cancer Trials Centre) and takes place in hospitals across the United Kingdom. A centralised database called MACRO is used for this purpose. Data collection and recruitment commenced April 2014. Recruitment completed at all sites on December 16, 2022 except at London and Manchester hospital sites where recruitment is due to complete March 31, 2022. A full list of all sites for recruitment and data collection can be found in the study protocol (see above).
	INSPIRE Trial: Patients recruitment and data collection was conducted at Princess Margaret Cancer Centre from March 21, 2016 to May 9, 2018. Data collection cutoff was July 18, 2019. Refer to Yang et al. 2021, https://doi.org/10.1038/s41467-021-25432
Outcomes	TRACERx Lung: disease-free survival is a pre-defined primary endpoint determined by RECIST v1.1 criteria. Last updated June 15, 2021.

INSPIRE Trial: Survival outcomes of the study was defined by progression free survival (PFS) and overall survival (OS). Clinical outcomes was determined by RECIST criteria and clinical benefit was defined as patients with complete or partial response or stable disease greater than 18 weeks. Refer to Yang et al. 2021, https://doi.org/10.1038/s41467-021-25432

Flow Cytometry

Plots

Confirm that:

- The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).
- The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers).
- All plots are contour plots with outliers or pseudocolor plots.
- X A numerical value for number of cells or percentage (with statistics) is provided.

Methodology

Sample preparation	Thawed samples were washed in sterile PBS to remove traces of DMSO and serum before staining with Zombie NIR viability dye (1:500 dilution in PBS for 15min/room temperature). Samples were then stained for lineage and differentiation markers for 15 minutes at 4oC (see key reagents table, all antibodies used at 1:100 dilution in FACS buffer), washed twice with sterile 4oC FACS buffer, kept on ice and immediately acquired on a BD LSRFortessa or sorted on a BD FACSAria Fusion.
	For in vitro activation assays, thawed samples were washed in sterile PBS to remove traces of DMSO and rested overnight in complete RPMI medium (10% FCS + penicillin/streptomycin) at 37oC/5% CO2. Rested cells were seeded at up to 200,000 cells/well in 200mcl of complete RPMI medium the following day. Cells received either no stimulation (complete RPMI medium only) or were stimulated with PMA (10ng/mI) and ionomycin (1mcg/mI). Brefeldin A (5mcg/mI) and anti-CD107A (1:400 final dilution) was added to all wells. Plates were centrifuged at 200g for 2 minutes and incubated at 37oC/5% CO2 for 5 hours. After 5 hours, cells were stained for surface lineage markers as described above. Following surface staining, samples were fixed in BD CellFIX and washed twice with permeabilization wash buffer (Biolegend) before staining for intracellular cytokines (1:100 final dilution of each antibody in permeabilization wash buffer) for 20 minutes at 4oC. Samples were then washed twice with permeabilization wash buffer and immediately acquired on a BD LSRFortessa.
Instrument	BD LSRFortessa or sorted on a BD FACSAria Fusion.
Software	FCS files were analysed using FlowJo version 10.
Cell population abundance	Between 50 and 500 cells were sorted for low input RNA sequencing as detailed in methods.
Gating strategy	Preliminary gates selected cells using an FSC and SSC gate. Doublets were excluded by plotting FSC area versus FSC height and excluding cells outside of a linear correlation. Live leukocytes were gated as CD45-high Zombie-NIR-low single cells. Subsequent gating was as described in supplementary data panels.

X Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.